Consciously Controlled Breathing Decreases the High-Frequency Component of Heart Rate Variability by Inhibiting Cardiac Parasympathetic Nerve Activity

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Heart rate variability (HRV), the beat-to-beat alterations in heart rate, comprises sympathetic and parasympathetic nerve activities of the heart. HRV analysis is used to quantify cardiac autonomic regulation. Since respiration could be a confounding factor in HRV evaluation, some studies recommend consciously controlled breathing to standardize the method. However, it remains unclear whether controlled breathing affects HRV measurement. We compared the effects of controlled breathing on HRV with those of spontaneous breathing. In 20 healthy volunteers, we measured respiratory frequency (f), tidal volume, and blood pressure (BP) and recorded electrocardiograms during spontaneous breathing (14.8 ± 0.7 breaths/min) and controlled breathing at 15 (0.25 Hz) and 6 (0.10 Hz) breaths/min. Compared to spontaneous breathing, controlled breathing at 0.25 Hz showed a higher heart rate and a lower high-frequency (HF) component, an index of parasympathetic nerve activity, although the f was the same. During controlled breathing at 0.10 Hz, the ratio of the low frequency (LF) to HF components (LF/HF), an index of sympathetic nerve activity, increased greatly and HF decreased, while heart rate and BP remained almost unchanged. Thus, controlled breathing at 0.25 Hz, which requires mental concentration, might inhibit parasympathetic nerve activity. During controlled breathing at 0.10 Hz, LF/HF increases because some HF subcomponents are synchronized with f and probably move into the LF band. This increment leads to misinterpretation of the true autonomic nervous regulation. We recommend that the respiratory pattern of participants should be evaluated before spectral HRV analysis to correctly understand changes in autonomic nervous regulation.

Keywords: autonomic nervous system; controlled breathing; heart rate variability; respiratory frequency; tidal volume

Introduction

Since the autonomic nervous system (ANS) plays an essential role in modulating physiological conditions during physical activity, mental stress, and even sleep (Hjortskov et al. 2004; Kuo and Yang 2009; Mendonca et al. 2009), understanding this system is key to providing proper care for patients with autonomic impairment and should be useful for elucidating abnormal regulatory mechanisms of the ANS in certain diseases.

The cardiac ANS comprises sympathetic and parasympathetic nervous activities. The relative contributions of these nervous activities to the heart underlie the beat-to-beat variability of R-R intervals (RRI), which is the elapsed time between two consecutive R waves on an electrocardiogram (ECG). Analysis of periodic fluctuations in heart rate, known as heart rate variability (HRV), has attracted considerable interest as a noninvasive method for quantifying alterations in cardiac autonomic regulation (Akselrod et al. 1981). Spectral analysis of HRV enables investigation of frequency-specific oscillations of heart rate fluctuation and helps decompose a series of consecutive RRI into a sum of sinusoidal functions of different amplitudes and frequencies (Moodithaya and Avadhany 2012). HRV mainly consists of two spectral components, one of low frequency (LF) and one of high frequency (HF), and it is widely accepted that the HF component of HRV reflects cardiac vagal nerve activity at the sinus node (Pomeranz et al. 1985; Berntson et al. 1993; Hayano et al. 1994; Task Force of the European Society of Cardiology the North American Society of Pacing Electrophysiology 1996). Pomeranz et al. (1985) showed atropine, a parasympathetic muscarinic blocker, practically abolished the HF component. Some studies demonstrated through spectral analysis that the HF component decreased in myocardial infarction (Lombardi et al. 1987), congestive heart failure (Axelrod et al. 1987), diabe-
Various pathological and physical conditions affect HRV by inducing changes in the cardiac ANS, although other factors, particularly respiratory parameters, may also affect HRV, since the HF component is associated with respiratory sinus arrhythmia (RSA), which is of respiratory origin (Novak et al. 1993; Brown et al. 1993; Pitzalis et al. 1998; Badra et al. 2001). RSA is a physiological phenomenon that is characterized by a decrease in RRI concurrent with inspiration when respiratory mechanisms in the brainstem attenuate the vagal efferent action to the heart and by an increase in RRI during expiration when the vagal efferent influence to the heart is activated (Tonhajzerova et al. 2013). Another mechanism regarding RSA is that a decrease in intrathoracic pressure that occurs during inspiratory phase causes a decrease in left ventricular stroke volume, resulting in a decrease in arterial blood pressure (BP). This decrease in BP probably leads to a decrease in RRI mediated by baroreceptor reflex (Larsen et al. 2010). Therefore, it is likely that an alteration in respiration is closely related to a change in HRV. In fact, it has been reported that the decrease in respiratory frequency ($f$) and increase in tidal volume ($V_T$) are positively correlated with an increase in the HF component, which suggests that the respiratory parameters themselves probably cause changes in HRV (Hirsch and Bishop 1981; Novak et al. 1993; Brown et al. 1993; Pöyhönen et al. 2004).

To avoid the respiratory impact on HRV, consciously controlled breathing has been used in experiments and clinical trials to standardize HRV measurements (Grossman et al. 1991; Piccirillo et al. 2004; Sin et al. 2010; DeBeck et al. 2010; Schaffer et al. 2013). However, it is possible that HRV is influenced by changes in $f$ and $V_T$ during the transition from spontaneous to consciously controlled breathing, which might lead to inaccurate estimations that deviate from the true autonomic nervous activity in the heart. In addition, the effect of consciously controlled breathing on HRV is not yet fully understood. Therefore, whether controlled breathing is required while assessing the ANS remains debated among researchers.

The aim of the present study was to clarify the effect of controlled breathing on HRV, including respiratory and circulatory parameters, and compare it with that of spontaneous breathing. We assumed that consciously controlled breathing altered HRV by inducing changes in respiratory parameters.

**Methods**

### Subjects

Twenty young healthy male volunteers aged 22.8 ± 0.7 years participated in our study. Their average height, body weight, and body mass index were 172.5 ± 1.3 cm, 62.4 ± 1.9 kg, and 20.9 ± 0.5 kg/m², respectively. All participants were non-smokers without any history of respiratory or cardiovascular disease. None of the participants routinely took medicine that affected autonomic nervous regulation or participated in sports at a competitive level. Written informed consent was obtained from all participants, and the study was approved by the Ethics Committee of the Tohoku University Graduate School of Medicine.

### Experimental protocol

Data were recorded in a quiet, air-conditioned laboratory at an ambient temperature of approximately 24°C and humidity of approximately 40% between 9:00 a.m. and 2:00 p.m. on a single day, considering the circadian rhythm of autonomic nervous activity (Bonnemeier et al. 2003). All volunteers were asked to refrain from drinking alcohol and caffeinated beverages for at least 12 h prior to data collection. They were also required to have a moderate amount of sleep the night before and not to eat for at least 2 h before the experiment.

The participants were seated in a comfortable chair and wore ECG electrodes and a face mask throughout the experiment. After a 5-min rest for stabilization, they were asked to consciously control their breathing to 6 and 15 breaths/min (0.10 and 0.25 Hz) for 5 min at random using a digital timer following spontaneous breathing without conscious control for 5 min. The breathing rate of 6 breaths/min was selected as it could induce dynamic changes in $f$ and $V_T$ during the transition from spontaneous to consciously controlled breathing and has been frequently used to investigate the influence of slow and deep breathing on the ANS (Hayano et al. 1994; Pitzalis et al. 1998; Ducla-Soares et al. 2007). The breathing rate of 15 breaths/min is close to the mean respiratory rate in adults and has been often used to standardize measurements obtained in spectral analysis of HRV (Brown et al. 1993; Bernardi et al. 2000; Piccirillo et al. 2004). Additionally, these consciously controlled breathing rates were included within the LF band between 0.04 and 0.15 Hz and the HF band between 0.15 and 0.40 Hz. All participants practiced and became accustomed to breathing at the required frequencies before the experiment (Pinna et al. 2006). Correct control of breathing was visually assessed according to the breath-by-breath $f$ displayed on the respiratory monitor.

Respiratory parameters, namely $f$ and $V_T$, were recorded continuously during the experiment by using a portable expired gas analyzer (AT-1100; Anima Co., Tokyo, Japan). Prior to data collection, the analyzer was calibrated using gas mixtures with accurately known concentrations of $O_2$ and $CO_2$. The validity and compatibility of the device are detailed in another report (Tajima and Ito 2006). The face mask was designed to fit comfortably without any air leakage and was fixed over the participant’s mouth. Systolic and diastolic blood pressure (SBP and DBP, respectively) was measured in the right upper arm by using a digital sphygmomanometer (CITIZEN CH-308B; Tokyo, Japan). The experimental procedure is depicted in Fig. 1.

ECG data were sampled at 1,000 Hz, digitized using an analog-to-digital converter, and stored on a personal computer for subsequent analysis. Spectral analysis was performed off-line by continuous wavelet transformation using Fluiclet (Dainippon Sumitomo Pharmaceutical Co., LTD., Osaka, Japan). LF and HF components were obtained as the area under the power spectral curve for the LF and the HF bands, respectively. In general, the HF component repre-
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Preceding text...


ds parasympathetic nerve activity at the sinus node (Pomeranz et

al. 1985; Berntson et al. 1993; Hayano et al. 1994; Task Force of the

European Society of Cardiology the North American Society of

Pacing Electrophysiology 1996), while the LF component probably

reflects both sympathetic and parasympathetic modulation of heart

rate (Pomeranz et al. 1985) and is strongly influenced by baroreflex

activity. In our study, to estimate the autonomic nervous activity in

the heart, we used the HF component as an index of cardiac vagal

nerve activity and the ratio of LF to HF components (LF/HF) as an

index of sympathetic nerve activity (Pagani et al. 1986).

Statistical analysis

All measurements including RRI, BP, respiratory parameters,
the HF component, and LF/HF, were averaged under each breathing
condition. Since no significant difference was observed between
the values under the first and second spontaneous breathing conditions,
we compared the values between the first spontaneous breathing
condition and each controlled breathing condition using the paired t-test.
A linear regression analysis was used to test the relationship between
changes in respiratory parameters and the HF component. Data are presented
as mean ± standard error of the mean. A value of p < 0.05 was considered
to be statistically significant.

Results

Respiratory rate

The participants’ breathing rate was 14.8 ± 0.7 breaths/min during spontaneous breathing without conscious control. During consciously controlled breathing at 0.25 and 0.10 Hz, they breathed almost correctly at 14.9 ± 0.1 and 6.4 ± 0.04 breaths/min, respectively. No significant difference was observed between spontaneous breathing and controlled breathing at 0.25 Hz.

Changes in tidal volume during controlled breathing

At both controlled breathing rates, VT was significantly higher than that during spontaneous breathing. During controlled breathing at 0.10 Hz, the increase in VT was reasonable because all participants breathed slowly during the transition from spontaneous to controlled breathing. However, an unexpected increase in VT was also observed during controlled breathing at 0.25 Hz, even though the mean respiratory rate was the same for spontaneous breathing and controlled breathing at 0.25 Hz (Table 1).

Circulation during spontaneous breathing and controlled breathing

No significant difference was found in the SBP and DBP between spontaneous breathing and either controlled breathing rate. However, the RRI were significantly lower during controlled breathing at 0.25 Hz than during spontaneous breathing (Table 1).

HRV during spontaneous breathing and controlled breathing

The HF component significantly decreased at both controlled breathing rates than during spontaneous breathing (Fig. 2). With regard to LF/HF, this was significantly lower during controlled breathing at 0.25 Hz than during spontaneous breathing. Conversely, LF/HF was considerably higher during controlled breathing at 0.10 Hz than at

Table 1. Comparison of VT, BP, and RRI between spontaneous breathing and controlled breathing at both rates.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Spontaneous breathing</th>
<th>Controlled breathing (0.25 Hz)</th>
<th>Controlled breathing (0.10 Hz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>VT (ml)</td>
<td>534.6 ± 23.4</td>
<td>594.7 ± 29.5*</td>
<td>1,030 ± 76.5**</td>
</tr>
<tr>
<td>SBP (mmHg)</td>
<td>113.6 ± 1.9</td>
<td>114.6 ± 1.5</td>
<td>110.5 ± 2.0</td>
</tr>
<tr>
<td>DBP (mmHg)</td>
<td>64.6 ± 1.3</td>
<td>65.7 ± 1.6</td>
<td>63.6 ± 1.6</td>
</tr>
<tr>
<td>RRI (msec)</td>
<td>857.8 ± 27.3</td>
<td>842.5 ± 29.4*</td>
<td>860.7 ± 23.5</td>
</tr>
</tbody>
</table>

Values are presented as mean ± standard error of the mean. *p < 0.05 and **p < 0.01, spontaneous breathing vs. controlled breathing at 0.25 Hz or 0.10 Hz using the paired t-test.

0.25 Hz = controlled breathing at 15 breaths/min, 0.10 Hz = controlled breathing at 6 breaths/min. VT, tidal volume; SBP, systolic blood pressure; DBP, diastolic blood pressure; RRI, R-R intervals.
spontaneous breathing (Fig. 3).

**Relationship between changes in respiratory parameters and the HF component during controlled breathing at 0.25 Hz**

Figs. 4 and 5 show how changes in the actual measurements of \( f \) and \( V_t \) affected the actual value of the HF component during the transition from spontaneous breathing to controlled breathing at 0.25 Hz. In subjects who increased their respiratory rate during controlled breathing at 0.25 Hz, the HF component decreased. Therefore, a significant inverse correlation exists between the change in respiratory rate and the HF component during the transition from spontaneous breathing to consciously controlled breathing at 0.25 Hz (Fig. 4). On the other hand, in subjects who increased their \( V_t \) during controlled breathing at 0.25 Hz, the HF component increased. Thus, the increase in \( V_t \) from spontaneous breathing to controlled breathing at 0.25 Hz was correlated positively with the increase in the HF component, although this correlation was not statistically significant (Fig. 5).

**Discussion**

The results of the present study suggest that both controlled breathing rates affect HRV by inducing the changes in respiration, since we found that controlled breathing at 0.25 Hz altered the HF component and LF/HF, while the controlled breathing at 0.10 Hz largely affected HRV, especially LF/HF.

**Spontaneous respiration vs. consciously controlled respiration**

During controlled breathing at 0.25 Hz, a reduction in the HF component and RRI was observed, suggesting that the conscious control of respiration leads to inhibition of vagal nerve activity in the heart, and subsequently, a decrease in RRI. The decrease in the HF component observed in this study is consistent with the findings of previous studies (Sloan et al. 1991; Patwardhan et al. 1995b; De Meersman et al. 1995). Patwardhan et al. (1995b) showed that the mean values of heart rate and BP increased, while the HF component of HRV decreased, when their participants adjusted conscious control of breathing to their own spontaneous breathing pattern. In the present study, controlled breathing at 0.25 Hz also caused a decline in RRI. There could be two possible explanations for this finding. The first is the changes in respiratory parameters brought about by controlled breathing, which could affect the cardiovascular system as hypocapnia due to conscious hyperventilation causes a decrease in RRI (Van De Borne et al. 2001; Debreczeni et al. 2009). In fact, we found that the \( V_t \) was significantly greater during controlled breathing than during spontaneous breathing, although the mean \( f \) was almost the same between both breathing conditions, and this finding supports the view that mild hyperventilation occurs during controlled breathing at 0.25 Hz. The other possible reason could be that cortical input accompanied by the action of consciously controlled respiration induces alterations in circulatory regulation (Han et al. 1997). Controlled respiration requires mental concentration, which could tend to reduce the HF component (Sloan et al. 1991), as seen in trials where mental stress produces an increase in heart rate and BP (Pagani et al. 1991; Yoshino and Matsuo 2005). Additionally, De Meersman et al. (1995) found that in their study, all subjects experienced discomfort and a significant loss of vagal tone through
intervention involving adjustment of breathing rate and $V_T$. Although other findings regarding the effect of controlled breathing on HRV have found no significant difference in the HF component between spontaneous and controlled breathing (Hayano et al. 1994; Patwardhan et al. 1995a; Bloomfield et al. 2001) or have found an increase in this component during controlled breathing compared to that during spontaneous breathing (Pagani et al. 1986; Driscoll and Dicicco 2000), the conflicting results may be attributed to different experimental procedures and parameters,
including the controlled breathing rate and/or $V_T$, position, technique of respiration modulation, and method of HRV analysis.

Changes in respiratory parameters also have an impact on the HF component in terms of the relationship between $f$ or $V_T$ and the component as well as the inhibition of parasympathetic nerve activity at the sinoatrial node during consciously controlled breathing at 0.25 Hz. An inverse relationship between breathing rate and the HF component has previously been reported (Hirsch and Bishop 1981; Brown et al. 1993; Hayano et al. 1994; De Meersman et al. 1995). In particular, in the present study, $f$ may have had a greater influence on the HF component than $V_T$ since no significant relationship was observed between $V_T$ and the HF component, and the correlation coefficient was greater for $f$ than $V_T$ (Figs. 4 and 5). These findings indicate that $V_T$ has little effect on HRV, which is supported by the fact that the effect of $V_T$ on fluctuations in RRI is negligible (Hirsch and Bishop 1981; Brown et al. 1993; Cooke et al. 1998).

In sum, the results regarding controlled breathing at 0.25 Hz indicate that controlled breathing with mental concentration may affect ANS function as well as respiratory and circulatory regulation, and the change in $f$ is likely an important confounding factor in the interpretation of HRV results.

**Respiratory frequency within the LF band**

During controlled breathing at 0.10 Hz, a reduction in the HF component of HRV and a large increase in LF/HF were observed, but these results may lead to the incorrect assumption of withdrawal of cardiac vagal nerve activity and activation of sympathetic nerve activity, since we showed that RRI and BP remained almost unchanged throughout the intervention. In general, provided that the above-mentioned autonomic alterations occur, the RRI decreases and BP increases. This interesting phenomenon coincides with the findings of earlier reports (Sanderson et al. 1996; Strano et al. 1998; Bernardi et al. 2000). Bernardi et al. (2000) observed that in each task accompanied with a decrease in RRI, namely, reading aloud, free talking and verbal arithmetic, the LF component increased and the HF component decreased, while the $f$ under these conditions not only decreased but also shifted into the LF band of HRV (0.04-0.15 Hz). Some HF subcomponents, namely, RSA, are synchronized with $f$ (Brown et al. 1993; Pitzalis et al. 1998) and probably move into the LF band, indicating that the increase in the LF component in some mental activities during slow respiration can be determined from the effects of slowed respiration and/or increased sympathetic nerve activity. Similarly, the slow breathing of some athletes influences the spectral indices of sympathetic activity (Strano et al. 1998). Considering the unchanged RRI and BP in our study and the findings of previous studies, the significant increase in LF/HF during controlled breathing at 0.10 Hz also suggests that $f$ shifts strongly into the LF band from the HF band, resulting in a relative increase in the LF component and decrease in the HF component.

Changes in the breathing pattern have been reported in several physiological situations, including exercise and sleep (Mador and Tobin 1991; Eisele et al. 1992). Moreover, RSA was found to differ depending on the spontaneous breathing period of each individual (Ben Lamine et al. 2004), since some individuality in respiration pattern exists among humans (Shea et al. 1987; Benchetrit 2000). Considering these facts, evaluation of HRV using spectral analysis necessitates assessment of the participant’s respiratory style. Unless the breathing pattern is determined in subjects breathing at 0.15 Hz or less within the LF band, it may not be possible to accurately determine the influence of the LF component on the overall findings.

**Limitation**

A major limitation of our research is the effects of the face mask. In the present study, the participants wore a face mask throughout the experiment, whereby ventilation could be directly measured, since the extent to which $V_T$ changes the HF component during controlled breathing has not been determined thus far. It has previously been reported that the use of a mouthpiece and a noseclip or a face mask directly affects measurement results as the $V_T$ increases because of dead space, although the effects on minute ventilation and $f$ are not as consistent (Weissman et al. 1984; Rameckers et al. 2007). In addition, Furutani et al. (1997) observed that HRV was affected by the dead space created by the use of a face mask. In contrast, our pilot study on six healthy volunteers showed that the use of a face mask had no statistically significant effect on HRV, RRI, or BP, compared to when a mask was not used. Nonetheless, considering the findings of several studies collectively, the possibility that direct fitting with a face mask may have had an influence on HRV and cardiovascular regulation in the present study cannot be ignored.

**Clinical implications**

Spectral analysis of HRV in the past has provided clinically valuable information. Fukusaki et al. (2000) reported aging was associated with decreased HRV, especially the HF component, indicating attenuated parasympathetic nerve activity in the heart. In patients with severe coronary artery disease and congestive heart failure, decreased HRV was recognized as ANS impairment (Casolo et al. 1995). A decline in HRV has also been shown to be an independent predictor of fatal arrhythmic events and sudden death due to congestive heart failure (Fauchier et al. 1999). Several reports have shown ANS dysfunction among patients with diabetes mellitus (Lindmark et al. 2003), hypertension (Mussalo et al. 2001), multiple system atrophy (Kitae et al. 2001), and Parkinson disease (Ziemssen and Reichmann 2010). Additionally, the prevalence of sleep-disordered breathing may increase with age to a prevalence of approximately 20% in older patients relative to 5% to 10% in
younger populations in The Sleep Heart Health Study (Young et al. 2002). Moreover, sleep-disordered breathing such as obstructive sleep apnea and central sleep apnea is common in patients with heart failure, and its occurrence has been reported to range from approximately 50% to 80% (Lanfranchi and Somers 2003). Further, a cohort study suggested that obstructive sleep apnea hypopnea syndrome occurred in up to 83% of patients with uncontrolled hypertension despite them taking three or more antihypertensive agents at the optimum doses (Logan et al. 2001). Sleep dysfunction including sleep apnea is also observed in patients with Parkinson disease (Diederich et al. 2005). Considering all these findings collectively, the results of our study indicate that alterations in respiratory pattern caused by sleep apnea affect HRV, and it is therefore very important to assess respiration in older patients and in patients with respiratory disturbance before evaluating autonomic regulation of the heart using spectral HRV analysis.

Conclusions

Our study involving 20 young healthy male participants showed that controlled breathing at 0.25 Hz changed not only HRV but also $V_T$ and RRI compared to those with spontaneous breathing without conscious control, suggesting that controlled breathing, which requires mental concentration, affects the ANS as well as respiratory and circulatory modulation. In contrast, controlled breathing at 0.10 Hz mostly affected LF/HF, but the RRI and BP remained almost unchanged; if only the increase in LF/HF is considered, this may be misinterpreted as activated sympathetic nerve activity. Further, our results indicate that respiratory frequency should be regarded as an important confounding factor in the interpretation of HRV findings. Despite the notion that R-R variability is not a simple marker of vagal nerve activity but instead depends on the complex interaction between vagal and sympathetic nerve activity and respiration, little attention has been paid to respiration in many previous studies on spectral analysis of HRV. Therefore, we recommend that the respiration pattern of each participant be evaluated before spectral analysis of HRV in order to correctly understand the changes in true autonomic nervous regulation.

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Conflict of Interest

The authors declare no conflict of interest.

References


Controlled Breathing Decreases the HF Component of HRV


